

N THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of:

Lynch et al.

Appl. No. 09/961,381

Filed: September 25, 2001

For:

Model for Neurodegenerative **Diseases Involving Amyloid**

Accumulation

Confirmation No. 7154

Art Unit:

1632

Examiner:

Crouch, D.

Atty. Docket: 1819.0040001/MAC/LBB

First Supplemental Information Disclosure Statement

Commissioner for Patents PO Box 1450 Alexandria, VA 22313-1450 Sir:

Listed on accompanying Form PTO-1449 are documents that may be considered material to the examination of this application, in compliance with the duty of disclosure requirements of 37 C.F.R. §§ 1.56, 1.97 and 1.98. The numbering on this First Supplemental Information Disclosure Statement is a continuation of the numbering in Applicants' Information Disclosure Statement filed electronically herewith, in connection with the above-captioned application. A copy of each document is also provided.

Where the publication date of a listed document does not provide a month of publication, the year of publication of the listed document is sufficiently earlier than the effective U.S. filing date and any foreign priority date so that the month of publication is not in issue. Applicants have listed publication dates on the attached PTO-1449 based on information presently available to the undersigned. However, the listed publication dates should not be construed as an admission that the information was actually published on the date indicated.

Applicants reserve the right to establish the patentability of the claimed invention over any of the information provided herewith, and/or to prove that this information may not be prior art, and/or to prove that this information may not be enabling for the teachings purportedly offered.

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This statement should not be construed as a representation that a search has been made, or

that information more material to the examination of the present patent application does not exist.

The Examiner is specifically requested not to rely solely on the material submitted herewith.

This First Supplemental Information Disclosure Statement is being filed more than three

months after the U.S. filing date and after the mailing date of the first Office Action on the merits,

but before the mailing date of a Final Rejection, or Notice of Allowance, or an action that otherwise

closes prosecution in the application.

This IDS is being filed as two submissions on the same day. One submission is electronic

and one submission is non-electronic. It is believed that only one fee under 37 C.F.R. §1.17(p) is

required to cover both submissions. That fee is included with this non-electronic submission.

Attached is our PTO-2038 Credit Card Payment Form in the amount of \$180.00 in payment of the

fee under 37 C.F.R. § 1.17(p).

It is respectfully requested that the Examiner initial and return a copy of the enclosed PTO-

1449, and indicate in the official file wrapper of this patent application that the documents have been

considered.

The U.S. Patent and Trademark Office is hereby authorized to charge any fee deficiency, or

credit any overpayment, to our Deposit Account No. 19-0036.

Respectfully submitted,

STERNE, KESSLER, GOLDSTEIN & FOX P.L.L.C.

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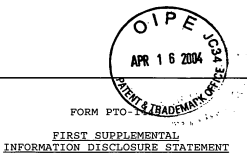
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APPLICATION NO. ATTY. DOCKET NO. 1819.0040001/MAC/LBB 09/961,381 APPLICANT Lynch et al. FILING DATE GROUP

FIRST SUPPLEMENTAL INFORMATION DISCLOSURE STATEMENT

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	AR	13	Ghiso, J., et	t al., "A 109 ursor protein	-amino-acid C-terminal contains a sequence, -18:1053-1059, Portland P	fragmen	t of Alzhein that promote	
	AS	13	Disease Tangl	insberg, S.D., et al., "Expression Profile of Transcripts in Alzheimer's isease Tangle-Bearing CAl Neurons," <i>Ann. Neurol. 48</i> :77-87, Lippincott Williams Wilkins (July 2000)				
	AT	<u>13</u>		n E-deficien	ory deficits and choline ot mice," <i>Neurosci. Lett</i>	. 199:1	-4, Elsevier	
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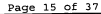


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	AR	14	Endocytosis a	at the Apical	Actin Microfilaments Pl but not the Basolatera 1 Biol. 120:695-710, Th	l Surfa	ce of Polari	.zed
	AS	14	Gouras, G.K., Pathol. 156:1	uras, G.K., et al., "Intraneuronal A $eta42$ Accumulation in Human Brain," Am. J. thol. 156:15-20, American Society for Investigative Pathology (January 2000)				
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	AR	<u>15</u>	Haass, C., et al., "Mutations Associated with a Locus for Familial Alzheimer's Disease Result in Alternative Processing of Amyloid β -Protein Precursor," J . Biol. Chem. 269:17741-17748, The American Society for Biochemistry and Molecular Biology (1994)					ırsor," <i>J</i> .
	AS	<u>15</u>	Haass, C., et al., "Targeting of cell-surface β -amyloid precursor protein to lysosomes: alternative processing into amyloid-bearing fragments," Nature 357:500-503, Macmillan Magazines (1992)					
	АТ	<u>15</u>	Alzheimer's D Consequences	Disease and S of the Time-	ry, Jr., P.T., "Models of Scrapie: Mechanistic Trut Dependent Solubility of aal Reviews Inc. (1997)	ths and	d Physiologic	cal
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	AR	<u>16</u>	1-3) on Amylo Hippocampal S	Harris-White, M.E., et al., "Effects of Transforming Growth Factor- β (Isoforms 1-3) on Amyloid- β Deposition, Inflammation, and Cell Targeting in Organotypic Hippocampal Slice Cultures," <i>J. Neurosci</i> 18:10366-10374, Society for Neuroscience (1998)					
	AS	<u>16</u>		Hoffman, K.B., et al., "β-Amyloid increases cathepsin D levels in hippocampus," Neurosci. Lett. 250:75-78, Elsevier Science Ireland (1998)					
	AT	<u>16</u>	Alzheimer's d	lisease mouse	que-independent disruptic models," <i>Proc. Natl. Ad</i> aces (March 1999)	on of ne	eural circui i. USA 96:32	ts in 28-3233,	
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	AR 17 Huang, TF., "What have snakes taught us about integrins," Cell. Mol. Life Sci. 54:527-540, Birkhäuser Verlag (1998)						fol. Life	
	AS	17	Immunoglobuli	Juse, W.D., et al., "Generation of a Large Combinatorial Library of the immunoglobulin Repertoire in Phage Lambda," <i>Science 246</i> :1275-1281, American association for the Advancement of Science (1989)				
	AT	<u>17</u>		ns by integri	n Nhieu, G., "Binding an n receptors," <i>Trends Mic</i>	crobiol.	. 2:10-14, E	
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	AR	<u>18</u>	pathway in br	ain parenchy	ification of the major A ma: Suppression leads to 143-150, Nature America	bioche	emical and p	athological
	AS	18	Janus, C., et Biophys. Acta	Janus, C., et al., "Transgenic mouse models of Alzheimer's disease," <i>Biochim</i> . Biophys. Acta 1502:63-75, Elsevier Science B.V. (July 2000)				
	AT	<u>18</u>	Kang, J., et resembles a c (1987)	al., "The pr	ecursor of Alzheimer's or receptor," <i>Nature 325:73</i>	disease 33-736,	amyloid A4 Macmillan J	protein ournals
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	AR	Kato, H., et al., "Graded expression of immunomolecules on activated microglia in the hippocampus following ischemic in a rat model of ischemic tolerance," Brain Res. 694:85-93, Elsevier Science B.V. (1995)							
	AS	<u>19</u>	Köhler, G., a antibody of p (1975)	Köhler, G., and Milstein, C., "Continuous cultures of fused cells secreting antibody of predefined specificity," <i>Nature 256</i> :495-497, Macmillan Journals (1975)					
	AT	<u>19</u>	Alzheimer Dis	sease Brains,	er-soluble Aß (N-40, N-4 " <i>J. Biol. Chem. 271</i> :40 ecular Biology (1996)				
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	AR	20	OTHER (Including Author, Title, Date, Pertinent Pages, etc.) Labat-Robert, J., "Cell-Matrix Interaction, Alterations with Aging, Involvement in Angiogenesis," Pathol. Biol. (Paris) 46:527-533, Expansion Scientifique Francaise (1998)					
	AS	<u>20</u>	Dermo-Epiderm	Le Varlet, B., et al., "Age-Related Functional and Structural Changes in Human Dermo-Epidermal Junction Components," <i>J. Investig. Dermatol. Symp. Proc.</i> 3:172 179, The Society for Investigative Dermatology, Inc. (1998)				
	AT	20	Marsh, M., an 285:215-220,	ıd McMahon, H American Ass	H.T., "The Structural Era sociation for the Advance	a of Endement of	locytosis," f Science (J	Science July 1999)
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	AS	<u>21</u>	Miller, D.L., et al., "Peptide Compositions of the Cerebrovascular and Senile Plaque Core Amyloid Deposits of Alzheimer's Disease," Arch. Biochem. Biophys. 301:41-52, Academic Press (1993)								
	AT	21	Morgan, D., e model of Alzh (December 200	neimer's dise	peptide vaccination preve case," <i>Nature 408</i> :982-985	ents mer	nory loss in re Publishir	ı an animal ıg Group			
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	AR	22	Morris, J.C., rules," Neuro	"The Clinic Dlogy 43:2412	al Dementia Rating (CDR -2414, Advanstar Commun): Curre	ent version s (1993)	and scoring	
	AS	Mucke, L., et al., "High-Level Neuronal Expression of Aβ ₁₋₄₂ in Wild-Type Human Amyloid Protein Precursor Transgenic Mice: Synaptotoxicity without Plaque Formation," <i>J. Neurosci.</i> 20:4050-4058, Society for Neuroscience (June 2000)							
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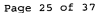
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	AS	<u>25</u>	Palmer, G.C., Neuropatholog (September 20	jies," Curr.	ction by NMDA Receptor A Drug Targets 2:241-271,	Antagon Benthar	ists in a Va π Science Pu	riety of ublsihers
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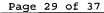
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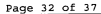
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